Coronavirus and Homo Sapiens

Pooja Natarajan¹ Muralidhar Kanchi¹ Vikneswaran Gunaseelan² Alben Sigamani² James Harmon³ Kumar Belani⁴

¹Department of Anaesthesiology and Critical Care, Narayana Institute of Cardiac Sciences, Narayana Hrudayalaya, Bangalore, Karnataka, India
²Department of Research, Narayana Health City, Narayana Hrudayalaya Limited, Bangalore, Karnataka, India
³Department of Surgery, University of Minnesota, Minneapolis, Minnesota, United States
⁴Department of Anesthesiology, University of Minnesota, Minneapolis, Minnesota, United States

Address for correspondence Muralidhar Kanchi, MD, FIAC, FICA, MBA, FASE, Department of Anaesthesiology and Critical Care, Narayana Institute of Cardiac Sciences, Narayana Hrudayalaya Health City, Bommasandra Industrial Area, Anekal taluk, Bangalore 560 099, Karnataka, India (e-mail: muralidhar.kanchi.dr@narayanahealth.org).

Abstract

The Spanish influenza pandemic of 1918 globally claimed death between 50 and 100 million lives. In India, it was referred to as “The Bombay Fever,” and accounted for a fifth of the global death toll at that time. The current outbreak of the novel coronavirus disease 2019 (COVID-19), a new human-infecting beta coronavirus, has demonstrated that the size of an organism does not reflect on its ability to affect almost an entire human population. COVID-19, first detected in December 2019 in Wuhan, China, spread rapidly worldwide. In humans, this disease ranged from flu-like symptoms to severe acute hypoxic respiratory failure. By appearance, this virus closely related to two bat-derived severe acute respiratory syndrome (SARS) coronaviruses. Although bats were likely the original host, animals sold at the Huanan seafood market in Wuhan might have been the intermediate host that enabled the emergence of the virus in humans. Under the electron microscope, the severe acute respiratory syndrome-coronavirus-2 (SARS-CoV-2) virus grips its receptor tighter than the virus behind the SARS outbreak in 2003 to 2004. The viral particle docks onto the angiotensin-converting enzyme 2 (ACE2) receptor and initiates viral entry. This review discusses the various aspects of the SARS-CoV-2 virus, its structure, pathophysiology, mechanism of interaction with human cells, virulence factors, and drug involved in the treatment of the disease.

Keywords
► antiviral treatment
► coronavirus spike protein
► cytokine storm

Introduction

A novel variety of coronavirus, the severe acute respiratory syndrome-coronavirus-2 (SARS-CoV-2) virus, reported late in 2019 from Wuhan, Central China, has now globally spread with ruthless speed.¹ It has expressed itself as a pandemic in our planet at all continents except Antarctica. The SARS-CoV-2 virus has been solely responsible and far exceeded the number of lives lost within a short period when compared to the 1919 pandemic and has had a significantly greater global economic impact.² The World Health Organization (WHO) informed that the coronavirus pandemic is “defining global health crisis of our time,” and has revealed the best and worst in humanity. On March 22, 2020, Mr. Narendra Modi, the Prime Minister of India, highlighted the scale of the challenge as follows: “Even World Wars I and II didn’t affect as many countries as the coronavirus has done.” SARS-CoV-2, the causative organism, is the name given to this virus, it is an enveloped RNA beta coronavirus that among other manifestations causes the respiratory disease called novel coronavirus disease 2019 (COVID-19; Fig. 1). Coronaviruses are a family of viruses, named for the crown-like effect created by spikes on their surface, which are surface proteins that help them invade human cells. Similar coronaviruses cause...
the common cold. What we’re dealing with now is a new, or novel coronavirus called SARS-CoV-2 that is effectively transmitted between humans and can cause a wide range of clinical conditions ranging from asymptomatic to a fatal infection in both adults and children.3

**Structure of Coronavirus**

Coronaviruses, a large family of viruses, have crown-like appearance due to protein spikes on their surface that help the virus to invade human cells. Coronaviruses belong to the family coronaviridae in the order nidovirales1,4 and are classified into four groups: alpha, beta, gamma, and the delta coronavirus. Alpha- and beta-coronaviruses infect mammals, gamma-coronaviruses infect avian species, and delta-coronaviruses infect both mammalian and avian species (Fig. 1). The examples of the alpha-coronaviruses include human coronavirus NL63 (HCoV-NL63), porcine transmissible gastroenteritis coronavirus (TGEV), porcine epidemic diarrhea coronavirus (PEDV), and porcine respiratory coronavirus (PRCV). Beta-coronaviruses include SARS-CoV-1 and -2, MERS-CoV (Middle East respiratory syndrome-coronavirus), bat coronavirus HKU4, mouse hepatitis coronavirus (MHV), bovine coronavirus (BCoV), and human coronavirus OC43. Gamma- and delta-coronaviruses include avian infectious bronchitis coronavirus (IBV) and porcine delta-coronavirus (PdCV). Coronaviruses are remarkably large, enveloped, positive-stranded RNA (ribosomal nucleic acid) viruses, and have the most voluminous genome, ranging from 27 to 32 kb.4 The nucleocapsid helical protein encloses the genome and is further surrounded by an envelope. The envelope has three structural proteins, namely, membrane (M), envelope (E), and spike (S) proteins (Fig. 2A). The M and E proteins facilitate viral assembly, whereas S protein is responsible for the viral attachment and entry into host cells. The S structural protein forms substantial protrusions from the virus surface, giving coronaviruses the look of having crowns (corona), is a key determinant of viral host response and is a major facilitator of the host-immune response.1

**The Life Cycle of SARS-CoV-2**

The invasion begins when the S protein binds to the host cellular ACE2 receptor. Binding triggers a conformation change in the S protein that leads to viral envelope fusion with the host cell membrane through the endosomal pathway. SARS-CoV-2 then releases RNA into the host cell (Fig. 3). Genome RNA is translated into viral replicating polyproteins pp1a and pp1ab that are then broken into smaller products by viral proteinases. The breakdown reaction produces a collection of subgenomic mitochondrial RNAs (mRNAs), by interrupted transcription and transformation into relevant viral proteins. Viral proteins and genomic RNA are collected into virions in the endoplasmic reticulum (ER), and golgi apparatus, and then transported via vesicles which are released from the infected host cell. The SARS-CoV2 S protein has three segments: a large ectodomain, a single-pass transmembrane anchor, and a short intracellular tail. A receptor-binding subunit S1 and a membrane-fusion subunit S2 form the ectodomain. Electron microscopy (EM) studies indicate that the spike is like a clove-shaped trimer with three S1 heads and a trimeric S2 stalk.6-11 When the virus enters the host cell, the S1 head binds to a receptor on the host cell surface, attaching the virus, enabling the S2 stalk to connect with the host and viral membranes, allowing the viral genomes into the host cells (Fig. 2B). These crucial initial steps during coronavirus infection, namely, receptor binding and membrane fusion are primary targets for human therapeutic interventions.

**Coronavirus and Receptor Binding**

Coronaviruses demonstrate a pattern of receptor recognition.10 The alpha-coronavirus HCoV-NL63 and the beta corona SARS-CoV viruses, both recognize a zinc peptidase in...
the ACE2 receptor. The SARS-CoV-2 gains access into the cells by binding to the ACE2 receptor. Two binding hot spots have been identified on human ACE2: ACE2 residues Lys31 and Lys353.11-18 These two hotspots contribute significantly to virus–receptor binding. The coronavirus exists in two distinct configurations: (1) prefusion trimeric spike containing three receptor-binding S1 heads and a trimeric membrane-fusion S2 stalk and (2) postfusion trimeric S2 which is a six-helix bundle with exposed fusion peptides. A variety of triggers regulate the transition of the spikes from the prefusion to the postfusion arrangements. Receptor attachment and membrane fusion are critical determinants of the host response and tissue response is characteristic of coronavirus infection.

As coronaviruses bind to the ACE2 receptor, it was proposed that ACE inhibitors (ACE-1) and angiotensin receptor blockers (ARBs) may be associated with increased severity of illness among COVID-19 patients due to ACE2 receptor upregulation.19,20 Researchers have also hypothesized that inhibition of the renin-angiotensin system and increased levels of ACE2 may have a protective effect in acute lung injury. In a recent study evaluating a cohort of 205 hospitalized patients with COVID-19 infection, although ACE-1 did not affect severity of illness, they did provide a beneficial effect. In this study, 37 (18%) were on ACE-1, and a serial logistic regression analysis confirmed a significant decrease in the primary endpoints of death or admission to the intensive care unit (ICU) in those on ACE-1 compared to patients not on ACE-1.21,22 The possible protective and treatment effectiveness of ARBs (losartan) is currently being evaluated in paired multicentered randomized double-blinded control clinical trials initiated at the University of Minnesota (ClinicalTrials.gov: NCT04311177 and NCT04312009).

The Cytokine Storm and Secondary Hemophagocytic Histiocytosis

A devastating, rapidly fatal cytokine storm may occur in coronavirus infections, resulting in secondary hemophagocytic histiocytosis (sHLH), characterized by multiorgan failure ( Fig. 4). This cytokine storm is responsible for much of the problems encountered by patients with the SARS-CoV-2 infection because the storm is a hyperimmune response to the virus resulting in a dysregulated, and accelerated expression of proinflammatory cytokines interleukin (IL)-2, IL-6, IL-8, and tumor necrosis factor (TNF). The predominant feature of sHLH includes reduced white blood cell counts, specifically lymphopenia, elevated serum ferritin levels, and severe acute respiratory distress syndrome (ARDS). High-ferritin levels are particularly dangerous because they suggest the presence of a significant hyperinflammatory response with their consequences on the lung.23,24 Thus, the treatment goals are to lower the consequences of the severe cytokine storm during severe COVID-19 pneumonia. Since entry of SARS-CoV-2 in cells is dependent on the connection of viral proteins S with cellular receptors and activation of viral proteins by proteases of host cells, this could be one area of virus inhibition. Thus, factors that affect the clathrin-mediated endocytosis (a procedure that is partly regulated by microtubules remodeling) could potentially decelerate viral infection of cells.25 Colchicine is another drug that could be helpful because it has high bioavailability in granulocytes and monocytes. Its property to bind unpolymerized tubulin heterodimers, to form a stable complex effectively inhibits microtubule dynamics and is a nonselective inhibitor of the NLRP3 inflammasome, a major pathway element in the development of ARDS.26 The options for cytokine inhibition include possible corticosteroid or intravenous immunoglobulin administration. A randomized
controlled trial with tocilizumab, an IL-6 blocker, is currently recruiting patients in China. Other treatment options are based on clinical findings and include heparin, serine protease inhibitors, such as ulinastatin, high-dose vitamin C, continuous renal replacement therapy (CRRT), and high-volume hemofiltration, as adjuncts in the care of critically ill patients with COVID-19 infection. Anti-IL-1 therapy with anakinra is being evaluated for patients with severe COVID-19 and shLH. Evidence for a severe cytokine storm and mortality include reports by Tu et al. reporting higher levels of IL-6, C-reactive protein (CRP), and D-dimer levels in nonsurvivors. These findings offer information regarding the characteristics of severe COVID-19 infection and support further investigation regarding the use of immunomodulators. An expert consensus from China recommended cytokine clearance using an artificial liver blood purification (ALBP) systems. Plasma exchange, plasma absorption, and hemofiltration or plasma filtration have also been considered as alternative therapies.

Coronavirus and Hemoglobin Metabolism

The pathological mechanism coronavirus causing COVID-19 remains enigmatic and mysterious. A report that looked at biochemical indices of 99 patients with COVID-19 demonstrated abnormal hemoglobin metabolism. This article revealed a decrease in the hemoglobin and neutrophils counts associated with elevated levels of serum ferritin, erythrocyte sedimentation rate (ESR), CRP, albumin, and lactate dehydrogenase (LDH).

Exclusive Molecular Mechanism Explaining COVID-19 Expression

COVID-19 may not resemble the usual types of ARDS as seen in our routine critical care practice. The key pathogenic molecular step of SARS-CoV-2 is to attack the 1-beta chain of hemoglobin, attacking the porphyrins, dissociating, and releasing iron into the circulation. The ORF1ab, ORF10, and ORF3a proteins of the virus attack the heme on the 1-beta chain of hemoglobin. The virus binds deoxygenated hemoglobin readily compared to oxygenated hemoglobin resulting in resistant hypoxia coupled with rapid multiorgan failures. Following hemolysis of red blood cells, viral proteins bind to the hemoglobin and the virus enters the host cell via the spike-CD147 pathway. The virus interferes with the heme anabolic pathway and causes the disease. The free iron released into the circulation is toxic, causing powerful oxidative damage to the lungs. Free iron toxicity results in inflammation of alveolar macrophages that leads to characteristic changes seen on computerized tomographic scans of the lungs. The host attempts to compensate by accelerating hemoglobin synthesis consistent with improving hemoglobin values noted in these patients. Another compensatory mechanism that addresses the iron load is the increased ferritin levels documented in these patients. One cause for monocytosis seen in these patients may be the need for additional macrophages to engulf the excess iron load. The cause of lymphopenia may be that during white blood cell differentiation, the monocytes line is favored rather than lymphocytes line. As the iron load and hemoglobin increases, blood viscosity is also increased. This in combination with the hypercoagulable state may cause diffuse micro- and macrocirculatory thrombosis and the markedly elevated D-dimer levels seen in these patients. Postmortem studies have demonstrated that the ARDS picture is misleading and that the primary pathologic process is disseminated thrombosis. Mechanical ventilation without addressing this issue may cause more lung damage. Early and aggressive anticoagulation can be life-saving in these patients. Chloroquine phosphate competes with porphyrin and binds to the viral protein, thereby inhibiting the viral protein’s attacking the heme and the binding to porphyrin.

Thrombogenesis and Coagulopathy in COVID-19

A fulminant coagulopathy is described in patients with COVID-19 pneumonia. The hypercoagulable state is created by endothelial dysfunction leading to excessive thrombogenesis and inhibition of fibrinolysis. Hypoxia is a noted trigger of the procoagulant pathway leading to venous thrombosis. Postmortem examination of patients who died following a critical illness due to COVID-19 demonstrated microthrombosis in the pulmonary vessels. In a study of 183 consecutive patients with COVID-19, D-dimer and fibrin degradation products were significantly elevated, and the prothrombin and partial thromboplastin times were higher among nonsurvivors as compared to survivors. Among nonsurvivors, 71.4% had evidence of disseminated intravascular coagulation compared to 0.6% among survivors. The incidence of thrombotic events, including acute pulmonary embolism, deep vein thrombosis, acute ischemic stroke, acute myocardial infarction, and arterial embolism was evaluated in 182 COVID-19 patients admitted to three hospitals in the Netherlands. Thrombotic events were noted in 31% of patients in this study, acute pulmonary embolism being the most common complication (82%). Increasing age and the presence of coagulopathy were independent predictors of thrombotic events. Clinically significant coagulopathy with the presence of antiphospholipid antibodies was reported among three patients with COVID-19. In a retrospective, observational study of COVID-19 patients, who received anticoagulant therapy with unfractionated or low molecular weight heparin, was contrasted with those that had no anticoagulant treatment. A sequela-induced coagulopathy score (SIC) was calculated based on platelet count, international normalized ratio (INR), and the sequential organ failure (SOFA) score. On multivariate logistic regression analysis, patients with an SIC score were treated with unfractionated or low molecular weight heparin had a significantly lower 28-day mortality compared to those who did not receive anticoagulant therapy. Thus, should anticoagulation be part of the therapy.
<table>
<thead>
<tr>
<th>Drugs</th>
<th>Mechanism of action</th>
<th>Status of the clinical drug trial</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antiviral</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Remdesivir</td>
<td>Antiviral interferes with virus RNA polymerases to inhibit virus replication</td>
<td>U.S. Food and Drug Administration permitted emergency use authorization on May 1, 2020. Clinical trials have initiated in India</td>
</tr>
<tr>
<td>Favipiravir</td>
<td>Antiviral inhibits viral RNA polymerase, thus interfering with viral replication</td>
<td>Two clinical trials have achieved its primary endpoint</td>
</tr>
<tr>
<td>Lopinavir/ritonavir</td>
<td>Antiviral protease inhibitors</td>
<td>Trials did not achieve its primary endpoint</td>
</tr>
<tr>
<td>Oseltamivir (Tamiflu)</td>
<td>Antiviral drug approved for treatment of influenza A and B. It targets the neuramidase distributed on the surface of the influenza virus to inhibit the spread of the virus</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Umifenovir (Arbidol)</td>
<td>Antiviral impedes trimerization of SARS-CoV-2 spike glycoprotein and inhibits host cell adhesion like that of influenza virus hemagglutinin</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>EIDD-2801</td>
<td>Antiviral incorporated during RNA synthesis and then drives mutagenesis, thus inhibiting viral replication</td>
<td>Prepared for trial</td>
</tr>
<tr>
<td>CD24Fc</td>
<td>Antiviral immunomodulator against inflammatory response</td>
<td>Under evaluation</td>
</tr>
<tr>
<td><strong>Antimalarial</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chloroquine/hydroxychloroquine</td>
<td>Antimalarial endosomal acidification fusion inhibitor anti-inflammatory activity</td>
<td>Reduction of COVID-19 virus load reported. Results from ongoing clinical trials awaited. Its recommendation for treatment has been withdrawn by multiple agencies, due to potential toxicity. Does not help postexposure51</td>
</tr>
<tr>
<td><strong>Anticoagulants</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heparin</td>
<td>Anticoagulants reverse the hypercoagulability in severe cases</td>
<td>Proven trial</td>
</tr>
<tr>
<td><strong>Anti-inflammatory</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>NSAIDs anti-inflammatory</td>
<td>Controversial: avoid if usual contraindications present</td>
</tr>
<tr>
<td>Colchicine</td>
<td>Anti-inflammatory used in gout Inhibitory effects on macrophages; for COVID-19 with cardiomyopathy shown to reduce inflammation in the cardiac myocytes</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Anakinra</td>
<td>Modified human IL-1 receptor antagonist (IL-1RA) used in Rheumatoid arthritis</td>
<td>Under evaluation</td>
</tr>
<tr>
<td><strong>Immunological</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tocilizumab and sarilumab</td>
<td>Humanized mAb targeting IL-6</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Bevacizumab</td>
<td>Humanized mAb targeting VEGF</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Baricitinib</td>
<td>attenuates proinflammatory response by inhibiting JAK and blocks virus entering host cells through inhibiting AAK1</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Lenzilumab</td>
<td>Humanized monoclonal antibody that targets CSF2/GM-CSF</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>IFNs</td>
<td>Immune enhancer inhibits viral RNA transcription, protein translation and posttranslational modification, thus suppress virus replication</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>NK cell therapy</td>
<td>Immune enhancer direct cytotoxicity and immunomodulatory capability</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>IVIg</td>
<td>Immune enhancer passive immunity and anti-inflammatory effects</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Drugs</td>
<td>Mechanism of action</td>
<td>Status of the clinical drug trial</td>
</tr>
<tr>
<td>-----------------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>Corticosteroids</td>
<td>Reduces proinflammatory cytokines and possess antifibrotic property</td>
<td>Low dose recommended</td>
</tr>
<tr>
<td>Cepharanthine/selamectin/mefloquine hydrochloride</td>
<td>Inhibit infection of simian Vero E6 cells with pangolin coronavirus, whose S protein shares 92.2% amino acid identity with that of SARS-CoV-2, prevents viral entry</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Antihelminthic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Niclosamide and ivermectin</td>
<td>Anthelmintic drug, virus replication inhibitor</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Nitazoxanide and tizoxanide</td>
<td>Suppress proinflammatory cytokines in PBMCs and IL-6 in vivo</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Alternative medicine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chinese medicines</td>
<td></td>
<td></td>
</tr>
<tr>
<td>LHQW</td>
<td>TCM prevention and treatment for influenzas</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Xuebijing injection</td>
<td>TCM endotoxin antagonist, anti-inflammatory agent and anti-coagulant is used for sepsis</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Antibacterial</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Azithromycin</td>
<td>Antibacterial proven to be active in vitro against Zika and Ebola viruses</td>
<td>Positive data for the use, along with hydroxychloroquine, in a COVID-19 clinical trial. Not recommended in combination with HCQ because of cardiac arrhythmias</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Inhibitors of TMPRSS2 serine protease</td>
<td>Cleavage and activation of the S protein of SARS-CoV that is required for membrane fusion and host cell entry is mediated by TMPRSS2</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>rhACE2</td>
<td>ACE2 blocker binds to virus S-protein, thus protects host lungs from virus attack</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>iNO</td>
<td>Vasodilator potent and selective pulmonary vasodilation and antimicrobial activity</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Camostat mesilate (Foipan)</td>
<td>Synthetic serine protease inhibitors were developed for the treatment of oral squamous cell carcinoma, dystrophic epidermolysis, exocrine pancreatic enzyme inhibition, and chronic pancreatitis</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Nafamostat mesilate (Buipel)</td>
<td>A synthetic serine protease inhibitor approved in Japan for the treatment of acute pancreatitis, DIC and for anticoagulation in extracorporeal circulation: It inhibits MERS-CoV S protein-mediated viral membrane fusion with TMPRSS2-expressing lung Calu-3 host cells by inhibiting TMPRSS2 protease activity</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Dapagliflozin</td>
<td>Used to treat type-2 diabetes and, with certain restrictions, type-1 diabetes, adults with heart failure with reduced ejection fraction to reduce the risk of cardiovascular death</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Convalescent plasma</td>
<td>Antiviral plasma from recovered patients provides protective antibody</td>
<td>Early trials showing promising results.</td>
</tr>
<tr>
<td>Supportive</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vitamin C</td>
<td>Boosts immunity by stimulating IFN production, supplying lymphocyte proliferation and enhancing neutrophil phagocytic capability</td>
<td>Under evaluation</td>
</tr>
</tbody>
</table>
Table 1 (Continued)

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Mechanism of action</th>
<th>Status of the clinical drug trial</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin D</td>
<td>induces secretion of antimicrobial peptides and has immunomodulatory property</td>
<td>Under evaluation</td>
</tr>
<tr>
<td>Zinc</td>
<td>necessary for the immune system and has antiviral activities</td>
<td>Under evaluation</td>
</tr>
</tbody>
</table>

Abbreviations: ACE2, angiotensin-converting enzyme 2; AAK1, adaptor associated protein kinase 1; COVID-19, novel coronavirus disease 2019; DIC, disseminated intravascular coagulation; GM-CSF, granulocyte-macrophage colony stimulating factor; HCQ, hydroxychloroquine; IFN, interferon; IL, interleukin; iNO, inhaled nitric oxide; IVig, IVIG=intravenous immunoglobulin; JAK, janus kinase; LHQW, Lianhua Qingwen; mAb, monoclonal antibody; MARS-CoV, Middle East respiratory syndrome-coronavirus; NSAIDs, nonsteroidal anti-inflammatory drugs; NK, natural killer; PBMCs, peripheral blood mononuclear cells; rhACE2, recombinant human angiotensin-converting enzyme 2; SARS-CoV-2, severe acute respiratory syndrome-coronavirus-2; S protein; spike protein; TCM, traditional Chinese medicine; TMPRSS2, transmembrane protease/serine subfamily member 2; VEGF, vascular endothelial growth factor.

of patients with COVID-19. In view of a high incidence of thrombotic complications among patients with COVID-19, the International Society of Thrombosis and Haemostasis (ISTH) recommended administration of prophylactic low molecular weight heparin to all hospitalized patients with COVID-19 in the absence of active bleeding ensuring that platelet counts are greater than 25,000/μL, regardless of the INR and activated partial thromboplastin time (APTT). This strategy is expected to reduce the incidence of a sepsis-like coagulopathy and prevent venous thromboembolism.40

Drugs Used for COVID-19 Treatment

According to the WHO, there are no available vaccines nor specific antiviral treatments for COVID-19. Care for patients with COVID-19 includes isolation, social distancing, hygiene, treatment of symptoms, supportive care, and institution of experimental protocols. On May 1, 2020, the United States gave Emergency Use Authorization to the antiviral remdesivir for people hospitalized with severe COVID-19.31 In March, WHO initiated the “SOLIDARITY Trial” to assess the treatment efficacy of four existing antiviral compounds which are favipiravir, remdesivir, lopinavir, and hydroxychloroquine (HCQ; or chloroquine [CQ]).50 On March 16, 2020, the first clinical trial of a vaccine started that consists of a harmless genetic code copied from the virus that causes the disease at Seattle, United States.42 The following are the drugs suggested from literature searches against coronaviral disease (see Table 1).

Hydroxychloroquine

There is contradictory evidence regarding the use of HCQ in COVID-19 infection. CQ, used in treating malarial and autoimmune diseases, also confers considerable broad-spectrum antiviral effects even against SARS-CoV-2. HCQ is chemically like CQ but with lower ocular toxicity and has proven to be efficacious in containing SARS–CoV-2 in vitro.52,53 CQ phosphate inhibits terminal phosphorylation of ACE2, and HCQ elevates the pH in endosomes, which participate in virus cell entry. While HCQ may have benefit, it appears that additional investigation is required before committing to the routine use of HCQ against COVID-1954

QT Interval (ECG) Prolongation with the Hydroxychloroquine–Azithromycin Combination

Both HCQ and azithromycin are known to prolong the QT interval. Thus, patients on this combination require close cardiac monitoring. While the combination has been used successfully in some reports with in vitro efficacy of the combination.56 Chorin and associates describe the occurrence of significant QT prolongation including the occurrence of torsade de pointes when such a combination is used in the treatment of COVID-19.57 Thus, regular monitoring is required, especially among patients with renal dysfunction.

Remdesivir

The nucleoside analogue remdesivir has in vitro activity against SARS-CoV-2.58 It was originally developed to treat the Ebolavirus disease. It is an adenosine analogue which inserts into viral RNA chains, causing the premature breaking of the chains.58 On May 1, 2020, the U.S. Food and Drug Administration granted Gilead Sciences, Inc. Emergency Use Authorization of remdesivir to be prescribed by licensed health care providers to treat adults and children hospitalized with severe COVID-19.59 Severe COVID-19 is defined as patients with an oxygen saturation (SpO₂) ≤ 94% on room air or requiring supplemental oxygen or requiring mechanical ventilation or requiring extracorporeal membrane oxygenation (ECMO) and a heart–lung bypass machine. It was administered on a compassionate basis to 61 patients with COVID-19 who had an oxygen saturation of less than 94% on room air or required supplemental oxygen. Remdesivir was administered intravenously in a dose of 200 mg on day 1, followed by 100 mg per day for 9 days. Clinical outcomes of 53 of the 61 patients were analyzed. At baseline, 30 patients (57%) were invasively ventilated and 4 patients were on ECMO. The median follow-up period was 18 days. The level of the oxygen support device (ECMO, invasive mechanical ventilation, noninvasive ventilation, high-flow oxygen, or low-flow oxygen) could be successfully weaned in 36 patients (68%). Seventeen of 30 patients (57%) who were invasively ventilated were successfully liberated from the ventilator.
extubated. Twenty-five patients (47%) had been discharged, and seven (13%) had died at the time of follow-up. The mortality among invasively ventilated patients was 18% (6/34); mortality was 5% (1/19) among those who did not receive invasive ventilation.60

Favipiravir

Favipiravir is an antiviral used against influenza.61 It is an oral pyrazine carboxamide derivative and guanine analogue developed by Toyama Chemical in Japan. Favipiravir selectively inhibits the RNA-dependent RNA polymerase of RNA viruses and induces lethal RNA transversion mutations, thereby producing a nonviable virus phenotype; this phenotype cannot bind to E2 glycoprotein and nucleocapsid and its binding energy to viral E protein, ORF7a, and ORF1ab is higher than porphyrin. The binding energy of E protein and favipiravir is more than 2,700 times the binding energy of porphyrin. The primary function of E protein is to help the virus enter host cells, which suggests that favipiravir acts by effectively preventing the viral entrance to human host cells. The WHO and the European Union has initiated clinical trials to test remdesivir, CQ and HCQ, lopinavir/ritonavir (LPV-r), and LPV-r plus interferon (IFN) β-1a in COVID-19 patients worldwide in the SOLIDARITY Trial and in the DisCoVeRy Trial.62-64

Lopinavir/Ritonavir with or without Interferon Beta-1A

LPV-r is a specific protease inhibitor, fixed-dose combination medication used for the treatment and prevention of human immunodeficiency virus (HIV). Interferon-beta-1a is a cytokine in the interferon family used to treat multiple sclerosis (MS), produced by mammalian cells. Concomitant use of ritonavir and lopinavir could increase the plasma half-life of lopinavir through cytochrome P450 inhibition in the liver. Kim et al evaluated triple combination therapy with LPV-r, ribavirin, and IFN and has shown clinical effectiveness for MERS.65 A randomized control trial (MIRACLE trial) was initiated to determine the therapeutic efficacy of LPV-r combined with interferon β-1b in patients infected with MERS-CoV.66 Studies observed that treatment with LPV-r compared to the standard care group was not associated with any change in time to clinical improvement, and mortality at 28 days was similar in both groups. Treatment with LPV-r did not reduce viral RNA load and SARS-CoV-2 RNA remained detectable at 28 days in 40.7% of the patients in the LPV-r cohort. However, patients in the LPV-r cohort demonstrated fewer complications, lessor need for invasive respiratory support, and had fewer secondary infections than did patients who did not receive LPV-r treatment.67

Passive Antibodies

The transfusion of convalescent plasma collected from patients who had recovered from COVID-19 to patients who were newly infected is being investigated in an active clinical trial initiated at Mayo Clinic (expanded access to convalescent plasma for the treatment of patients with COVID-19. ClinicalTrials.gov identifier: NCT04338360). Transferring purified and concentrated antibodies via transfusion of convalescent plasma is a method of passive immunization. Neutralization of the virus is expected with this therapy, antibody-dependent cellular cytotoxicity and subsequent phagocytosis may be possible.68

Steroids

Steroid administration has shown to benefit patients in the acute phase of the disease.69 The WHO does not currently recommend corticosteroid therapy in other viral diseases, such as for patients with dengue. One concern is that the glucocorticoid-mediated stimulation of the hypothalamic-pituitary-adrenal axis may drive lymphocytopenia or promote exaggerated proinflammatory responses which eventually worsen the pathogenic condition.69,70 According to the surviving sepsis guidelines, there was a reduced length of stay in the ICU and in turn reduced cost, with the use of low dose steroids.71,72

Vaccine Trials

A number of phase-1 vaccine trials are already underway.73-79

Conclusion

COVID-19 has been the infection of the century and has surprised clinicians and scientists with its structure and mechanism of infecting host cells, mimicking ARDS, and enveloping hemoglobin resulting in severe hypoxia with an intense immune response and multiorgan failure. The molecular docking technology identified the binding site of viral proteins to porphyrin. The virus infects cells with ACE2 receptors; the immune cells produce antibodies leading to immune-mediated hemolysis. The elevated levels of inflammation result in a cytokine storm, causing multiple organ failure. Lung damage occurs with thrombosis and leads to hypoxia. Various drug trials are ongoing to find the ultimate cure for this viral disease. Careful selection out of this extensive menu of drugs available, with adequate knowledge about their side effects and, most importantly, prevention being the ultimate “mantra” in the treatment of COVID-19.

Mr. Sigesh AK, Department of Marketing and Development, Narayana Health, Bangalore, Karnataka, India, for helping us with the editing of diagrams.

Conflict of Interest
None declared.

Acknowledgment
The authors would like to thank Mr. Pavan Krishna Kanchi, MS, Chemical engineering, PhD scholar, IIT, Guwahati, Assam, India, for helping in making the first draft of the manuscript.

References
with the editing of diagrams.

Narayana Health, Bangalore, Karnataka, India, for helping us

hypoxia. Various drug trials are ongoing to find the ultimate

failure. Lung damage occurs with thrombosis and leads to

immune-mediated hemolysis. The elevated levels of inflam-

receptors; the immune cells produce antibodies leading to

viral proteins to porphyrin. The virus infects cells with ACE2

docking technology identified the binding site of

enveloping hemoglobin resulting in severe hypoxia with

mechanism of infecting host cells, mimicking ARDS, and

surprised clinicians and scientists with its structure and

COVID-19 has been the infection of the century and has

acute phase of the disease.69  The WHO does not currently re-

Steroid administration has shown to benefit patients in the

Steroids

sequent phagocytosis may be possible. 68

therapy, antibody-dependent cellular cytotoxicity and sub-

convalescent plasma for the treatment of patients with

much has changed. Available at: https://www.downtoearth.

nerlogists? Why and to what extent? The emerging

P

Nephron 2020;144(5):253–254

exerts a distinctive strategy for interacting with the ACE2

Br


sys

2006;80(14):6794–6800


Zheng M, Song L. Novel antibody epitopes dominate the antigenicity of spike glycoprotein in SARS-CoV-2 compared to SARS-CoV. Cell Mol Immunol 2020;17(5):536–538


36 COVID-19 disease: ORF8 and surface glycoprotein inhibit heme metabolism by binding to porphyrin. Available at: https://chemrxiv.org/articles/COVID19_Disease_ORF8_and_Surface_Glycoprotein_Inhibit_Heme_Metabolism_by_Binding_to_Porphyrin/11938173/3. Accessed August 10, 2020


COVID-19 and human beings  Natarajan et al.


50 WHO launches global megatrial of the four most promising coronavirus treatments. Available at: https://www.who.int/news-room/detail/2020-03-03-who-launches-global-megatrial-of-the-four-most-promising-coronavirus-treatments.

Accessed August 10, 2020


76 Caddy S. Developing a vaccine for covid-19. BMJ 2020;369:m1790

